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Polyurethane resins in water dilutable basecoats having low flash and quick-drying characteristics.

Tie) This investion relates to the field of polyumethane costings for use in automobile basecoutthercost systems. pile particlas, it is investion relates to the discovery that incorporating a long-yid-an entreview (and of a least 50%. Only weight of the carbonyle soil component used to make polyester restricts which are butter incorporated into provide the provides basecost composition entallising for temperature statch carbonations. These for propurature fasts characteristics are sethibled even where the basecost is deposited at 50-90% relative

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Polyurethane Resins in Weter Dilutable Basecosts Having Low Flash And Quick-Drying Characteristics

BACKGROUND OF THE INVENTION

Multi-layer systems have been utilized to cost automobies for a number of years, but the early sidevelopment of these systems necessarily employed organic schents. As environmental regulations became more stringent, and the cost of organic solvents trose, organic-bonne baseoust systems became less destrable. The recent research emphasis in the area of multi-layer systems, especially baseoust systems have been destrabled. The recent research emphasis in the area of multi-layer costores.

The shift from organic solvents to water for dispersing and applying resins in multi-layer systems solved many of the environmental and cost problems associated with the use of organic solvents. Water-borne systems however, here resulted in other problems.

The application of a multi-layer coaling to an automobile body, for entiriple, would be greatly feoritative by a system that provides for quick-right off entire multi-map and set the application of a coatery as quick-right of the coatery of the provides of the coatery of the coatery of the provides of the coatery of the

efficiency.

The present invention is directed to polyurethane coatings to be used in formulating basecoat compositions of multi-layer coatings systems. The nested of this invention are shown to possess the sequentials of being quick-daying outing and after application. Purthermore, the resens of this invention also enablest superfor coating characteristics, for example, opportunity of earth of the property of

ment. Roation, and life effect of the metallic pigments in the paint film. When non-metallic pigments are used, the resists of the present invention exhibit excellent foccasine effect.

This invention, therefore, relates to the field of polyurature coatings for use in automobile, be associately-easing systems, in particular, this invention relates to the discovery that increaseding either continued to the control of the coating of the carbonylic action of a feast 50% by weight of the carbonylic and component used to make polyester regists which are justified into polyvershare energy provide basedoct compositions existed.

low temperature flash characteristics. These few temperature flash characteristics are exhibited even when the basecasts are deposited at 50-90% relative humbly.

The polyrenthane restin, produced by the reaction of the above-described polyester restin and a polystocyantee mixture, although useful as a cessing composition for a number of bubstrates is especially useful as a basecular for administration. Coalings containing polyrenthane synetysists with an automatical production of the coalings containing polyrenthane synetysists with an activities with a supervision of the production of the coalings containing polyrenthane synetysists with an activities of the coalings contained to the coaling containing polyrenthane synetysists with an activities of the coaling containing polyrenthane synetysists with an activities of the coaling containing polyrenthane synetysists with an activities of the coaling containing polyrenthane synetysists with a containing containing the coaling containing polyrenthane synetysists with a coaling composition of the activities of the coaling containing polyrenthane synetysists with a coaling composition of a number of the coaling containing polyrenthane synetysists with a coaling containing polyrenthane synetysists with a coaling containing polyrenthane synetysists with a coaling containing polyrenthane synetysists and coaling containing polyrenthane synetysists with a coaling containing polyrenthane synetysists and coaling containing polyrenthane synetysists with a coaling containing polyrenthane synetysists and coaling containing polyrenthane synetysists with a coaling containing polyrenthane synetysists.

long chain latty acid comprising at least about 50% of the acid component in the polyester resin have shown to be particularly useful for water-home bascoest compositions used in multi-leyer systems. It is an object of this invention to provide polyure-there resins that can be incorporated into basecoat formulations to provide low fish and quitt-formy characteristics.

It is an additional object of this invention to provide polyester resins which can provide favorable low flash and quick-drying characteristics to polyurethane resins.

It is a further object of this invention to provide water-borne basecoat compositions having favorable coating and cosmetic characteristics and additionally provide for manufacturing efficiencies which result from the low-flash, quick-drying characteristics.

It is also an object of this invention to provide a method of producing the resins and basecoat compositions described herein.

It is a further object of this invention to provide a method for coating a metallic or plastic substrate so utilizing the resins and basecost formulations of the present invention.

These and other objects of the present invention are furthered by incorporating polyurethane resins into basecost formulations.

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SUMMARY OF THE INVENTION

automobile coating system.

The polyurethane redins are comprised of, in part, polyecter resins formed from a carboxylic acid component and a alcohol having at least 2 hydroxyl moeities. Specifically, this invention relates to an s anionic polyurethane coating composition comprised of:

1. A polyester component produced by condensing extraorytic acid component with alcohols having at least 2 hydroxy moeities wherein the cathorytic acid component is comprised or at least about 50°, weight of a long chain hydrophobic carboxytic acid contenting compound having between 18 and 80 carbon storns; and

2. A miscare of a compound having at least 2 isocyante groups, a multi-functional compound having at least the active hydrogen functionality and at least one cathorylic and terrocounty and optionally, a compound having at least the active hydrogen groups, for example, olds, difficial, distrinst, or compound having at least the active hydrogen groups, the optional compounds active district and having miscare of these active hydrogen groups, the optional compounds active district and the miscare to produce a copylumetrate resist containing the cathorylic acti groups. The free cathorylic acting groups he enterliated to produce a water-objective polymentum exist.

The polyurethane resin described above can be formulated as a water-dispersed basecoal resin along with a gried resin, a cross-linking agent, thicktopic or inacting common agents, bicketers, pigments, aluminum andro mice particles, bushfiling agents, water-flows, sunficarias, stabilizers, postations, wetting egents, departing agents, adhesion promoters, debarrers, catalysts, and additional polymers, for example 20 a branch-basin polyester among other ingordents.

After fermulation, the basecoalt compositions can be sprayed or electrostically deposited onto the authoritie body, preferably, in one or two coats. Generally, two even coats of basecoalt are applied with a one immunity fields between coats. After deposition of the basecoalt before application of a high solid content clear coat, it is generally preferred to Teach about 60% of the water from the basecoalt for optimum 35 spossarous and to eliminate washer boil of the clearcoat.

A preferred ambodiment of the vister-dispersible serioric real in relates to a polyvertheral product vibrating the polyverbares is formed with a mitter of an excess of discourants, a mail-functional composition and a fixed one active hydrogen functionality and at least one actively and of functionality, and a fixed premiserable objects reals. This inhister produces a usefulne-containing real intermediate history gives one to be fire inocyrates groups are polymer chain. In expectally preferred embodiments, the fire inocyrates groups are polymer chain. In expectally preferred embodiments, the fire inocyrates groups are polymer chain. In expectally preferred embodiments, the first one and the control of t

preferably, two or more. The polysection component is preferably formed from an alcohol component having at least about 2 hydroxy groups per miceloid job/yoh and a sathorpic and component. The carboxylic acid component is component to a set set about 5% by whigh (of a long polisin elsewher) and component is all of carbon and one of the carbon, the controlled component is at all of carbon and one of the carbon, the component is at all of carbon and the carbon and the

recognised but low flash and quick-drying characteristics of the basecost composition of this invention are ble neutral to having a high precenting of highly hydropticing groups in the polycent resinc. Clis to Coll carbonylis acid present a rings of compounds heigh subtable hydropholicity. Most preferably, this long chain carbonylis acid a disconsort case duel most greaterly as Call activative and from an a safety call and controlled and component preferably a distillational controlled and controll

polysetin ratio. The polyseter restins described hereinabove are select on virtually any elastomeric substants and are particularly useful when formistated into polyverance costings and used in basecus formistations for expectation onto material or plastic substants, expectally submorbly bodies. The opportune sensit or polysethan restant or plastic substants, expectally submorbly bodies. The opportune control polysethan restant or plastic substants, expectally submorbly bodies. The first control temperature polysethan restant, between them confidence softs as a function of a polysethan restant control and po The composition of the carboxylic acid component and polyor component employed to synthesize the polyester resists is such as to provide an excess of the polyor over and above the total number of equivalents of acid present in the misture. In other words, the resculant should be selected, and the stoichlorestic proportions of the respective solid and polyor components be adjusted to give hydroxy-terminated, polyerar molecules each theoretically, having a hydroxylindoxylind or 30 miles.

As stated above, the and mixture employing in forming the polyseter intermediate most preferably contains a S_S discharge/size did not receive as dimer and. Processes for forming this acid are well known and from the subject of numerous U.S. patient including Not. 2.482.751, 2.793.202 in 2793.201 and 2.895.121 or alternatively dimer fatty acid can be purchased from a chemical supply house (Empol 1010, a valuable from Emany Chamical Sur

O₂ dimer latity add fraction consists assentially of dimer (O₂ distribuyible adds) logisties with amounts up to about 30-25° of civilimer. Powers, note of still in the and rate to this dimerationer misture as "dimer" and the practice is followed hareless. The preferred grade contains 5°° dimer and 5°° timer. These opening products can be used in the form in which they are recovered from 1s° the polymerization unit. or they can be given a puril'al or complete hydrogenation trainment to restaurate to incommend to make the polymerization unit. Or they can be given a puril'al or complete hydrogenation trainment to additional contractions and the second can be the second to find the polymerization at the missing contraction of the missing characteristics.

The polyurethanes of the present invention are advantageously storage stable and are, of course, water dispersible. The water dispersible is controlled by the amount of free carboxylic acid contained in the final restin particles, and the number of sall groups termed from those tree acid groups.

Coating compositions produced using the polyurethane resins described herein have exhibited low flash and quick drying characteristics surprising for a water-dispersible resin.

DETAILED DESCRIPTION OF THE INVENTION

The present invention netales to a vater soluble aimoric polyurethane resin produced by reacting an polysterior compress of the seat 50% by weight of the catelogic loci dominants of a containing compound, a multivariant containing compound compounds compound compositions, an additional compound compressing a compound having at least their hydrogen containing moleties. The resulting polyurethane intermediate has terminal incorporate groups or active hydrogen-containing moleties, depending upon the succlonariesy of the colorisation of the containing and containing moleties.

An aspecially preferred embodiment of the polyurethree restrict of the present invention relates to the formation of a urethrane product in which the intermedate polyurethrane ream described above has free isocyarate groups at the terminal positions of the polyurethrane reals. The isocyarate groups are then capped with an excess of a polytunctional alcohol having at least 2 alcohol groups, and prefereby at least 3 alcohol groups.

The acid component of the polyester is, of course, critical to the invention and is comprised of a mixture of all less should 59% by weight of a long child componity acid component therein previews 88 and 80 celson atoms. Preterably, the long chair carbonylic acid is a dischoulkit acid and most preferably, the clicatorylic acid is a Cy. dismes clicitorarylic acid of miner acid. Where the long chair carbonylic acid comprises less than 100% of the carbonylic acid component, the carbonylic acid component is also comprised for the or more stort-chairse carbonylic acid.

Preferably, the long chain faily acid comprises between about 50 and 50% by weight of the acid component of the polyeeter polyce, the periodical series improvedeble the long chain faily, and component comprises about 78-60% of the long chain faily, and component and in the grind sean, the polyeeter rean comprises about 576-50% by weight of the objecter series. Otherwise, the inspire the percentage of long crisis comprises about 576-50% by weight of the objecter series. Otherwise, the significant percentage of long crisis followers, the abstrategious fash-off characteristics must be basiness with miss effect that fine change in the carbonytic acid component has not in emallatic effects, dareability and other characteristics of the ways. including, in the case of grind resin, the ability to accomodate pigment.

The shorter chain cateroyle add component is comprised of a none-, dir or ligiter interfocusity, catebookle add or a moture of these activoryle sadd heliog cannot chain of 12 or whee catebookle add the component of the componen

The polyester resints are synthesized from the shove-described carboxylic acid component and an access of a polyel component. An excess of polyel is used so that the polyester resin preferably contains terminal hydroxyl groups. The polyel compounds preferably have an everage hydroxy-functionality of at

The polysater resis in most cases is comprised of one or more polyois, preferably a diol. Up to about 25 percent by weight of the polyol component may be a polyol having three or more hydroxy groups per molecule. Where polyols having three or more hydroxy groups are chosen, the result is a branched

molecule. Where polyois having three or more hydroxy groups are chosen, the result is a branched polyester.

While it is not always desirable to have a triol or higher multi-functional alcohol present because of the

With it is not expected to be a controlled to the controlled to th

26 than how. The dids which are usually employed in making the polyecter relate include allysiene glycots, such as ethylene glycot, propyiene glycot, bezivene glycot, and reagonary glycot. (3 havened) and other glycots such as bydogreader bighinghoid. Only the glycots are such as bydogreader bighinghoid. Only the glycots are such as propriet by the glycots are propriet by the glycots and the glycots. The glycots are glycots and the glycots are glycots and glycots. The glycots are glycots are glycots are glycots are glycots. In a glycots are glycots are glycots are glycots. In a glycots are glycots are glycots. In a glycots are glyc

as higher molecular weight polytiols.

The low molecular weight dols which are preferred in the instant invention are known in the art. They have hydrony values of 200 or above, usually within the range of 200 or 200. Such materials involved as alphated color, aparticularly advises polytes containing from 2 to 18 control aborts. Examples involve ethylene glycol, 1,4-bustenedict, cyclosolysatic dols such as 1,2-cycloheanedict and cycloheaned climited such as proceeding preferred cold in 1,6 Pleaseaned.

The resulting polysetter rest in preferably processed with direct fathy acid as the long chain carbonylic acid (soperative acid as the minor short-chain carbonylic acid component component and an excess of acid to the carbonylic acid component component and are seened as the short acid acid to that the resulting objecter polyel register is size between about 200 or 200 grams per equivalent of hydroxyl. Preferably, the polyester rest has a range between 700 and 900 grams per equivalent of the droxyl and most preferably, has about 700 grams per equivalent of hydroxyl.

To produce the polymethane reshs which are useful in basecost competitions of the present invention, the above-described polyester polyol is restated with a miniture of a polyisocyprate, a multi-functional or compound hering at least one active hydrogen group and at least one active-viric acid group, and optionally, a component comprising a chemical compound having at least two active hydrogen groups, but no earthoutile acid mouns.

The polyetter, polyecycurets and multi-functional compound may see the reacted in the same pot, or may be recast expunsible, copredict poor the celefier results. Securetial reaction procures see which are more ordered in structure. Both the polyester and multi-functional compound may serve as chain estenders to build up the polyverster buckforch structure, reaction of hydroxyl groups. However, to function at a chain estender, the multi-functional compound must have all least five active hydrogen groups. Where the multi-functional compound has only one extremely result in the second process of the result is chain termination. Additional chain extenders being at least the active hydrogen groups. Where the multi-functional compound has only one extremely result in the process of the polyvershame. Additional chain extenders being at least the active hydrogen groups but not controlled and groups may be active to invinces the cell healing of the chainst determination of the polyvershame. The process of the polyvershame controlled and the process of the process o

The organic polytecopante which is reached with the polyhydric material as described is essentially any polytecopante and is printerly a discopantes a.e., hydrocathon discopantes are substituted hydrocathon discopantes. Many such organic discopantes are shrown in the art. including polytecylare discopantes, hothered discopantes, 3.3-dimethyl-4.4-bitolographies, blotted discopantes, 3.3-dimethyl-4.4-bitolographies, blotted discopantes, 3.2-dimethyl-blotten-1.6 discopantes, 1.4-tramethylene discopantes, 1.6-discopantes, blottened discopantes, 4.5-discopantes, discopantes, discopantes,

While disocyanates are preferred, other multi-functional isocyanates may be utilized. Examples are 1,2,4-benzene trifsocyanate and polymetrylene polyphenyl isocyanate.

It is preferred to employ an alignatic dilaccyantas, since it has been found that these provide better rs color stability in the finished coating. Examples include 1,3-hearamethylene cliscoyantate, 1-bebyriene discoyantate, methylene bis (4-cyclotheny) (socyanate) and isophorone discoyanate. Mixtures of discoyanates can albo be employed.

The proportions of the discorpants, polyester, and multi-functional compound are chosen so as to provide an inexpresse terminated intermediate polyurethane resin. This can be accomplished by utilizing a so stochometric excess of polyteocynates. i.e., more than one isocyanate group per nucleophilic moiety (reactive with locycantel) in the other components.

For purposes of promoting water-solubility it is important to build acid groups into the polyurethane. For example, the presence of acid groups is capable of rendering the composition water-dilutable.

The acids that are employed to provide free acid groups in the polyurelarar reams of this invention are readily available. They contain at least one active hydrogen group and at least one carbonylic acid functionally. The acids by hydrogen group may be a thiol, a hydroxyl or an armine, with primary armines being considered to have one active hydrogen group. Examples of such compounds include hydroxyl carboxylic acids, amino acids, in the acids, armine acids, the acids, armined acids, armine acids, and hydroxylical acids.

Compounds containing at least 2 hydroxyl groups and at least one carboxyle acid are preferred. They or can be prepared from an aldehyde that contains at least two hydrogens in the alpha position. Such atteryides are reacted in the presence of a loase catalyst with two equivalents of formaldehyde to form a 2.2-hydroxymdriyl aidehyde. The aidehyde is then gently catalast of the acid by known procedures. The acids that are employed in the invention can be expressinged in simplication by Formula for the control of the control of the present of the control of the control of the control of the present of the control of the contro

wherein R represents hydroxymethyl, hydrogen, or alkyl of up to 20 carbon atoms and preferably up to 8 carbon atoms.

Specific likatrative examples of such acids that are employed in the invention include 2.2-dilydroxymetrly) acets acid; 2.2-dilydroxymetrly) acets acid; 2.2-dilydroxymetrly) poster acid; 2.2-dilydroxymetrly) perinancic acid, and the like. The preferred acid is 2.2-dilydroxymetrly presion acid.

Lorgar-chan polyurature restra can be dozimed by chain extending the polyurature chain with a compound or midme of compounds containing at least two acids hydrogen groups but hiving no carbonities deld group. For example diols, difficult, distaining source having a minuter of hydroxyl-thill, and maning oppus, for example, allocationines, ammost offers, for purposes of his aspect of the invention both primary and secondary amen groups are considered as harmly one active hydrogen. Availablation, the manages, enterprises or of effective minutes are considered as harmly one active hydrogen. Availablation, the manages, enterprises or of effective minutes are supported as the secondary and active the secondary preferred did in expecting 500,000 cross. The same dold used to synthesize the polytester component of the polytesteries ensire can be utilized here as well, White polyhydroxy compounds containing at least three hydroxy oppose are by to used as for the active flower.

produces branched polyuesthane resists. For purposes of the present invention, it is preferred to minimize the amount of branching in the polyurathane resis. Therefore, if polyhydroxy compounds are used, they are preferably limited to a very minor component of the polyurethane producing mixture. These higher functional polyhydroxy compounds include, for example, trimethylopropene, trimethylopropene, a rido, among other compounds.

more, Talmay cores compositions, and the desired in early manner using these compounds having at least. The polyveretime resin say be chain extended in early manner using these compounds having at least on the minister of polytocyptests, polyvester and multi-functional compound, or alternatively, may resect at an intermediate stope, to link two free scorvantsed crouses that are exceeded at the terminal endo of an intermediate observableme resid.

t la generally preferred that an intermediate polyuetitane resis produced by resolting the polyyeques resid and the minuser of polytocytes, multimortional compound containing at least 2 shrivery pages and one carbonytic acid group, and clash estender be terminated with the locoyanist, groups. To excomplish this, an excess of the polytocytestes component is used. Of course, his motion ratio of the other components will be adjusted according to the deleted characteristics of the Intermediate and final is odiviruethmer explains. The polytocytest components on more than about 50% by wheeling the polytocytest components on more than about 50% by wheeling and contained the polytocytest components.

reaction ministers and it is preferred that the opposite component comprises from about 20% to about 70% by weight of reactants in the minister.

In one especially desirable embodiment of the invention, a multi-functional alcohol is used to terminate the reaction (cap the feel socyanate groups) at the desired stage (determined by the viscosity and

as incoparate groups present), thereby also combinding residual hydronyl groups. Particularly destrain for such purposes are ambinacionolis, such as effanciamine, diethanolamine and the like, since the amino groups preferentially react with the isocyanate groups present. Multi-funccional alcoholis, such as ethylene glycol, trinethyliopropene and hydrony-terminated polysetters, can also be employed in this manner. White the ratios of the components of the polysects, the multi-functional isocyanate and the terminating the properties of the properties of the polysects, the multi-functional isocyanate and the terminating the properties of the properties of the polysects of the multi-functional isocyanate and the terminating the properties of the properties of the properties of the polysects.

White the ratios of the components of the polyester, the multi-functional isocyanata and the terminating a spent can be varied, it will be noted by those skilled in the art that the amounts should be chosen so as to excid gelletion and to produce an ungelled, unethane reaction product containing hydroxyl groups. The hydroxyl value of the unethane reaction product should be at least 5 and preferably about 20 to about 200. The amount of convision-smalled used in the midster is creditably beforem about 20% and 30% by well-off.

of the reaction in fire mixture, but will very depending upon the polyvener used, the add number of the first polyvenerth resize, with the determination of the first polyvenerth resize, and the determination of the first polyvenerth resize, the moreunt of polytecyanate will also very depending upon whether it is desired to have the intermediate polyvenerth resize intermination with the isolocystate force on with hydrony groups. Thus, where it is preferred to terminate the residence polyvenerth resize in the late of the polytecyanate force of the polytecyanate resize to use of the polytecyanate resize to the pol

The amount of multi-nuclional component having at least one active hydrogen group and at least one activoryle acid group abor may very depending upon the desired acid number of the final polyurethere resir. The final polyurethere resin that final polyurethere resin that final polyurethere resin that final polyurethere resin that and about 2th by upplied of the reactures of section of the multi-nucleral component comprises between about 1% and about 2th by upplied of the reactures of all of the chairs estimate, for example compounds having the above hydrogens but no carbonyle groups). It is preferrable that the acid number be highter, because as the acid number increases, the example interest that the acid number is the state of the polyurethere resin potentially increases. The practical upper limit of add number is that which negatively effects the for final registroof profession of the final area. Of course, the upper limit of add number is that or each of course, the upper limit of the cold number will very depending upon the chemical composition of the final polyurethere resin but and outlines that is upper limit of the old in the practical final profession of the practical during the size of the final profession of the practical during the size of the final profession of the practical during the size upper limit of the old in the practical during the size upper limit of the old is in present. He practical

The amount of chain extender, when used producing the polyurethane resin, varies between about 2% and 25% by weight of the reactants. The amount used will depend upon the amount of chain extension of desired and the desired aspect as of a convertation emiscuse.

After the polyverbance resin is synthesized, the fine cuboxylic acid groups are nevel-table with base of orm sail groups. Profeselyb, the base is an amino containing compound. Testing writes are generally preferred over primary and secondary antites because of the tendency of the primary and secondary antities to resident the simpostation conscribing agents. Preferred testing varies include this silpointaines, for se example, trimethyl and trieflysimine. Also preferred is trieflysnotamine. Particularly preferred is trieflysnotamine.

The polyurethane resins of the present invention are formulated, along with other components, into water dispersible basecoat compositions which are sprayed or electrostatically deposited onto metal or

plasts statuted in example, automobile bodies, in general, a polyverbine reini formulate de described the plant of the pl

As indicated, an signation dispersion of the polyverbane resin is solited as the principal or major venicle resin. In general, the principal or major venicle resin in general, the principal or major venicle resides in general, the principal or major venicle resides present in the basecoat composition. This preference polyverbane resin is a resin produced from a polyperar synthetised from other large size plantified polyperar principal princip

The polyvertaine reaction product as described above is mixed with an ammogist resm. Ammogist resm are advanced conferention products of melitame, var. and emiliar compounds. Products obtained from the seaction of formation/de with melitames, uses or benoguearment are mest common and are semantic and accordance of the season of the seas

While the aldehylde employed is most often formaldehyde, other similar condensation products can be a made from other sidehydes, for example, acetaldehyde, crotoneldehyde acrolein, benzeidehyde, furfural, and others.

This amino-attentive condensation product contain methylol or similer alivelia groups, and in most instance at least a protine of these aliquid groups are otherwised by a reaction with an actional to provide organic solvent-equilibrium and actional composition and actional composition and actional composition actional composition and actional composition actional composition action actional composition actional composition action actional composition action action actional composition action actional composition action actions actional composition action actions actional actional composition action actions actional actions actionate actions actionated actions actions actions actionated actions actionated actions actionated actions actions actions actions actions actions actions actionated actions actions

A gird main is also used in the basecoat composition of the present invention. While the payment are main may be compressed of a number of water soluble problemeler relient, at a present of the time remains a composition of the present of the time problement of the time remains a composition of the present of the time remains a composition of the present of the section of the se

Coult and previously criminals action should be used as a grief resist in embodiments of this invention in A preference districtly polyteches resist for use as a grief resist in embodiments of this invention in the country of th

Figments may be incorporated into the basecest to provide the destined commit characteristics. This is done by mixing pigments with the above-described pigment resin and optionally, with other additives to form a pigment paste. Any standard pigment known in the art may be used with rearis of the present invention so long as these pigments can be formulated without affecting the desired low fitsh and quickdriving characteristics.

50 Specific examples of the dye staffs or pigments may be inorganic or organic. for example, grachinc, curbon black, time chromats, stendson chromats, basic unchromate, lead conversate, lead conversate lead conversate lead of the conversate lead of

The preferred metallic pigments are metal powders preferably mixed with aluminum metal flakes. The preferred aluminum flate pigments are swillable from Silverine Corp. Lundord, Pennsylvania or the Educat Werks, Quintenstabl. West Garmany. The aluminum flate pigments provide the coasing with an enhanced "metallic venees". In a preferred embodiment of the present invention standard grade aluminum stabilization with probapties seets in suck Other metallic files pigments; for evanging, silver may size be used but these are usually prohibitive in cost and inferior in appearance. The metallic pigments may also be mixed with non-metallic pigments, but these are to be carefully chosen so as not to diminish the desired metallic office.

The regins used in the basecost are dispersed in delottated water, it is preferred that the delottated water has conclusione sending or less than 1.3 microthers* and most preletably less than about 5 microthers* to greened passing caused by the reaction of aluminum with water. Delottated water is about 50 microthers* to greened passing caused by the reaction of aluminum with water. Delottated water is about one on those to be analyzed with the decirace water. An especially preferred colvent is 8-bbs (Calsockev's Winth add midding, formulating and dispersing pupils of the bits beared middle of the color of the bits of the color of the color of the bits of the color of the bits of the color of the c

The organic abhent, which comprises at most about 80% of the bissecont composition, and preferably comprises about 10% to 20% by weight of the bissecont composition (including weel) may be selected to promote the dispensibility of individual components in the final bissecont composition (plasticizer characterisfics).

A rheology control agent is also preferably incorporated into the basecoat composition. The rheology control agent controls the viscosity of the resulting composition and is incorporated in amounts that will prevent sagging or running after a basecost is spreyed onto e verticel surface such as an automobile body. The direct result of incorporating a rheology control agent is to provide flow control, body and sprayability. Other favorable results of adding a rheology control agent are to enhance the flip effect of metallic flake cirments. In denosit a thicker coating and to achieve complete coverege of a substrate. The sprayed coatings containing these agents also exhibit greater operation of the metallic flake pigments on the final coated substrate. Rheology control agents which can be used in embodiments of the present invention include the furned silica compounds and the bentorite clays. Preferred furned silica compounds are the hydrophobic silica compounds, for example Aerosil R972, available from DeGussa Corporation. (Frankfurt. West Germany). Another rheology control agent which may be used, and in certain basecoat compositions, may be preferred is a synthetic sodium lithium magnesium silicate hectorite clay. An example of one such clay is Laconite RD, available from Laconte, Inc (Saddlebrook, New Jersey), in certain preferred embodiments rhectory control agents are mixed. The rhectory control agent when it is included, generally comprises about 0.1 to about 20 percent by weight of the basecoat composition and preferably comprises 30 between about 1 percent and about 5 percent by weight of the final basecoat composition.

in general, the particle size of the rheology control agent plays a role in the overall thiototopic properties of these restine. Rheology control agents in embodiments of this investion are suspended in the meetral. It may be proposed that the rheology control agents are suspended and function, at least in part, through coalcombine or electrostatic interactions.

as In general, the particle sizes can be from less than 0.1 microns to over about 200 microns. These sizes can be adapted to develop in part the rheology properties sought. In appropriate circumstances, the particle sizes may be from about 0.01 to about 10 microns.

in addition to a principal resist or major verifice install rand a grind resist, preferred basecast compositions also are compressed of all least about 5th by verigit of the resinous verhels of a harmched-chain projecter is added for improved application properties and improved polystation properties and improved projected from carbonic projected in the improved projected proje

Any additional agent used, for example, surfazants, fillers, stabilizers, wotting agents, dispersing gaints, addesien premoters, etc. may be incorporated into the basecost composition. While the agents are well-known in the prior art, the amount used must be carefully controlled to avoid adversely affecting the schooling and epish-princ characteristics.

in formulating the basecost compositions of the present invention, the order of addition of the individual components is often very important. As a rule, the cross-linking agent in a solvent is added to the rheology control agent in solution and thoroughly mixed. Thereafter, the major vehicle resh dispersion (neutralized

with amine) is added to the rheology control solution under agitation. If desired a slurry of aluminum metal flakes end/or mica particles (mica particles are used alone in the case where an aluminum metaltic veneer is not desired) in Butyl CellosolveTM is mixed with a premixed starry of a tranched-chain polyeeter resin and dimethylethanciamine. This mixture of aluminum is then aplitated with the sturry containing macroust whicle, cross-linking agent, and rheology control agent. Pigment gastes comprised of colourethane resin pigment, fillers, stabilizers, plasticizers and other additives are then mixed under agitation with the aboveresulting mixture. Pigment paste particles are prepared in a sand mill, attritor or other common milling equipment prior to use.

The pigment pastes may be prepared by mixing the aminoplast resin with about 1.4 of the total colvurethane resin to be added to the pigment paste. Pigment is added to this sturry under agritation for about 1/2 hour. The rest of the polyurethane resin is then added and the resulting paste is mixed for another helf-hour. The pH and viscosity of the paste is checked and any adjustments are made by adding delonized water and/or tertiary amine. The weight ratio of pigment to binder usually ranges between 0.15-5.0. The amount of plament ranges between 6 and 90% of the total weight of plament plus binder. Other 15 well-known methods of formulating prepared prepaint passes may also be used.

The final basecost composition is adjusted to a pH of 7.6-7.8 with a tertiary amine, for example, Nethylmorpholine. Viscosity may be adjusted using delonized water. Final basecoat compositions are comprised of the following components in the indicated weight ratios.

Table I

Ingredient	Amount (% by weight of of Solids of Final Basecoat composition)
Polyurethane resin	20-80%
Melamine	5-50%
Rheology Control Agent	0-20%
Branched chain Polyester	0-35%
Pigment	2-65%

20

The basecost compositions described bereinshove can be applied to the metal or plastic substrate in one or two costs using for example an air atomizer (Rinks Model RI) spray our available from Rinks Manufacturing Corporation, Franklin Park, Illinois), or by using other conventional spraying means, The basecoat compositions may also be applied electrostatically. The basecoat compositions are preferably sprayed at 50-80psi, and a relative humidity of between 50 and 90% (optimally et 60-80% relative humidity) and a temperature of 70-90° F.

After being deposited, the basecost compositions are flash dried within a temperature range of about room temperature to about 145 degrees F for between 30 seconds and about 10 minutes using warm air blowing at a relative humidity of 5-40%. The preferred flash temperature is about 120 degrees F which is carried out for preferably between about 1 and 5 minutes. The flash conditions described herein result in about 90-95% of the solvents (water plus organics) being flashed from the basecoat in this short period of

After the first basecost is deposited, a second basecost can be deposited over the first without drying iffash off), or alternatively, a clearcoat may be deposited over the Eashed basecoat. Any number of clearcoat compositions known in the art may be used. Any known unpigmented or other transparently plamented coaling agent is in principle, suitable for use as a clearcoat. A typical top coat composition

contains 30-70% film forming resin and 30-70% volatile organic solvent. After the clear coat is coated onto the basecoat layer, the multi-layer coating is then baked to cross-link the polymeric vehicle and to drive the small amount of residual water and organic solvent from the multilayered polymeric composition. A preferred baking step involves heating the coated substrate for a period 55 of 10-60 minutes at a temperature of between 150 and 300 degrees F. The baking step cures the coating to

The invention will be further described in connection with several examples which follow. These examples are shown by way of illustration of the invention and are not meant to limit the scope of the

invention. All parts and percentages in the examples are by weight unless otherwise indicated

Polyurethane Example 1

Preparation of a Polyurethane Resin

- 10 A polyester polyol risen in prepared by charging a maction vessel (flask with a factionating column) with 551 B (15% of the polyester resinf of toesthates can: 1820 g, 164-59) Empol 1010 (client year) and in establish from Emery Chemical Col., and 1055: 1g, 162-59; in 13-Requested and 100 g, of tolware. Additional tolusione may be added to fifth the 58 pc. The financiary size for the concondensation was removed. Outing his heating 2837; g, of water was distilled off. Heating associated to the concordinate of the control of the control
- toluene is then vacuum stripped at 200° C to produce a polyvester resin for use in the polyvesthere resin.

 If this point, 897.9 g, of the above-synthesized polyvester resin 43.0 g, of dirently/oi propionic acid, 18.1 g, of neopenylighyod, 243.0 g, of lose) discorpanate and 300 g, of methyl sobouty featone are charged to a the reactor and heated at reflux (about 120 degrees C) until a constant isocyanate value is obtained, so 3.8 g, or thireactively process is then added to the reactor and the batch is alleved to reflux for an
- 30 36.8 g. of trimethylol programs is then added to the reactor and the batch is allowed to reflux for additional one hour. At this point, it is relingen purple is timed off and the batch is coulded to 8 degree C. 28.8 g. of climethyletherolamine and 100 be of water is then added using a protion of the water as a rinse. The batch is the male indexed to at utilit becomes homogeneous (bibut 5 minuses) and then 2048.71 g. of water is added over 2.00 minuse period under vigorous agistion.
 At the end of this addition the motives is distilled on high heat with vigorous agistion to remove water.
- and methyl isobutyl ketons. The water is then returned to the batch and the approximately 300 grams of methyl isobutyl ketone which was distilled off is discarded. 236 g. of rebutsoil is added and the batch is held at 80 degrees C for 30 millious. The batch is then dropped and filtered through a 10 million filter to give a polyester-unstainar vehicle for use in the basecost composition of the invention. The resulting of disneration but a solidis control of 30% and Gardon viscositiv of 25.

Polyurethane Dispersion 2

Preparation of Polyester Resin

A reaction vessel is charged with 1995 g. of adjois audi, 1995 g. of dimer audi, and 2490 g. of 1.6 hexaredid, and 139 g. toliumen. The mixture is heated under integen to 209 degrees C, removing where until an acid number less than 8 is reached. Remaining toluene is vacuum stripped to produce a polyester resin having solidac context greater than 95%.

4s Polyurethane Dispersion Preparation

857.4 g. of the atomic polyseter is mixed with 14.6 g. neopenty (pylout, St.) i.g. dimethylopiroptonic acts, 305.5 g. (supphorno difficoupsants, 7.1 g. methyl ethyl selaton, and 250.5 g. methyl ethyl isotone are refluend until a constant isotoprates value is obtained. At this point, 24.5 g. of detherodamine is at sided and be mixture as leaf of 20 mixture, 24.5 g. of direttly-plantomine and 11.6 g. decistand where the 25 g. isotopry abortal are at sided and all-week to make the first directly and appropriate the size of 250 g. decistand where the directly and appropriate directly application. The existing dispersion has a cubic constant of 25% and an appropriate Outdoore viscosity.

Polyurethane Dispersion 3

Preparation of Polyester

770 g, dimer acid, 230 g, 1,6-hexanediol, and 25 g, toluene are charged and the resulting mixture heated to 200 degrees C. Heating is continued, removing water, until an acid number less than 10 is achieved. The remaining toluene is then removed under vacuum.

Polyurethane

70 g. polysatir above, 128 g. reopenfyl glycol, 43 g. dimethylolyropionic acid, 244 g. isophorone dissocyantas, 77.8 g. methyl othyl kations, and 185.3 methyl armyl kation are macted using the procedure for polyurethrane disportion. 2. The resulting dispersion has a solids content of 20% and a Gardner viscosity of 71.

Branched Polyester 1

2594 g. of dimer acid. 2594 g. of 1.8-houseadol, and 744 g. of isophibalic acid are chapped and the misture heated in 16 flageres C under integrow with agilistic until acid marker of 10 or less is reached. 10 The insizure is then cooled to 150 degrees C and 1000 g. of timellist unity-riske is added slowly, and refunded until an add marker of 2002 is reached. After occing to 150 degree C or less. 729 g. of busy Colicohoeth and 1459 g. of housed are added. The resulting polyester has a solids content of 70% and a Sardher riske table VII-VI.

Branched Polyester 2

1330 g. dimer acid and 799.5 g. 1.8 hazinedick, we charged and heated to 196 degrees C under ritrogen with adjustion, Heating is continued until an acid number less than 10 is reached. The mixture is 30 than cooled to 190 degrees C and 434.1 g. timealfic anhydide is added slowly and heated until the acid number falls below 30, 335 g. buyly glycol and 570 g. ri-batterol set time added with agriation. The requiring polysester solition has a solide content of 170% and 5 deriver viscosing 10 deriver viscosing 10 deriver.

35 Branched Polyester 3

888.7 g. 1,5 hexanoidol. 1348.2 g. dimer acid, and 386 g. incontinuit and are heated at 195 degrees C until an acid number less than 8 is achieved. 208.6 g. incoellife, anhydride is then added slowly under agitation and heat applied until an acid number less than 30 is achieved. A 21 minute of n-butanoi and 40 butly (slycol are then added, until 70% solids is reached. The resulting branched polyester resin had a Gardner rescent of U.

PREPARATION OF COATING AGENTS

The composition of the coating agents is shown in Table 2, where the numbers denote parts by weight. The following notes refer to components listed thereof:

so Thickener 1:

Paste of Aerosil R972 (Degussa) hydrophobic furned silica sand milled with appropriate polyurethane grind resin and melamine in water, organic solvent mixture at 11% strength.

Thickener 2:

Pasts of synthetic sodium lithium magnesium silicate hectorite clay, Laponite RD (Laponte), 2% strength in delonized water; the paste is prepared by stirring with Cowles blade in water for one hour.

Thickener 3:

Paste of Laponite RD 3% strength in delonized water. Prepare as Thickener 2.

Titanium Dioxide Pigment Paste:

41 % concentration of DuPont R-960 titanium dioxide sandmilled with appropriate polyurethane grind rs resin and melamine.

Melamine Besin:

Commercially available methanol-etherfled melamine/formaldehyde resin, solids content 90% by weight in n-butanol.

Aluminum Pfoment I:

Silberline SS-5251 AR post treated with 4.5% Vircopet 40 (phosphate ester commercially available from Albright & Wilson, Richmond, Va.)

ao Aluminum Ploment II:

Stapa Hydrolac WH-R607 from Eckart Worke

Aluminum Pigment III:

Stapa Hydrolac WH-8487 from Eckart Werke

Thickener 2	40	37		37	38					_
Thickener 3			25					13	13	
Melamine Resin	4	4	4	4	8	7	4	1	2	
Butyl Cellosolve	1	1	1	1	1	2			1	
Polyurethane 1	40			40					18	
(30%NV)										
Polyurethane 2		44			42		38	17		
(26%NV)										
Polyurethane 3			44			38				
(26%NV)										
Aluminum I (54%NV)	6			6		6	6			
Aluminum II (65%NV)	5								
Aluminum III (65%N	W)		5		6					
Butyl Cellosolve	1	1	1		4	2	1			
Polyester 1 (70%)	6						6	3		
Polyester 2 (70%)			6	6						
Polyester 3 (70%)		6								
Dimethylethanolami	ne 2	2	2	2	1	6	6	1		
5% Strength in Wa	ter									
Thickener 1						27	27	17	17	
Titanium Dioxide P	aste							48	49	
Deionized Water			12	4		12	12			

EXAMPLES 1 to 4

The melamine restin and Busyl Celisopole are premised and acted to the hickers under agilation. The polyure/state dispersion is then acted to this mixture under agitation. An aluminum sturry is made by first is mixing the aluminum pipment and busyl celisosive, then acting the polyester restin, and then finally persecutating this sturry with the 5% DMEA solution. The aluminum sturry is then acted to the polyure/share/stickers/melamine institute under agitation.

re Example 5

Half of the measurine restin and buyli Cellosohell's age premixed and added to the thickener under agistion. The polyurethere desposation is the national, an elementary large inseparably by mixing a summirum plgment, remaining buyli Cellosohell's and measurine restin. The aluminum sturry is then added under significant to the rest of the paint. The pill is then adjusted with 5th disnet/benchamine in water.

Example 6

20 An aluminum slumy it made with aluminum pigment, melamine resin, and Butyl Cellosolve under agitation. The polyurethane dispersion is added to the aluminum slumy. Thickner is then added under aditation, off is adjusted with 5% DMEA and viscosity is adjusted with desortized water.

as Example 7

The polyurethane dispersion, melamine resin, and butyl Cellosolve are mixed with agitation. An aluminum sturry is made as in Example 1 to 5 and added to the first mixture under agitation. The thicker is added under cigitation. Viscosity is adjusted with delonized water.

Examples 8 and 9

Matamine resin and Butyl Cellosolve are premixed and added under agilation to Thickener 3. The polyurethane dispersion is bit hen added under agitation. In Example 8, the polyster rosin is preneturalized with 5% DMEA and then added under agitation. Thickener 1 (18972 paste) is added but when the tittanium dioutide paste is added both under agitation.

The invention has been described in detail with particular reference to preferred embodiments thereof, but it will be understood that variations and modifications can be affected within the spirit and scope of the invention and that the scope of the invention is to be determined by the claims appended hereto.

Claims

- An anionic polyurethane resin for use in a multi-layer costing system comprised of the reaction product of:
- (a) a polyester component with an atchde having at least two hydroxyl groups wheeler said carboxylic acid component is comprised of at least about 50% by weight of at least not long-plain carboxylic acid having between 18 and 60 carbon atoms and at most about 50% of at least one short chain of disproporties.
 - (b) a multi-functional compound having at least one active hydrogen group and at least one carboxylic acid functionality;
- (c) a compound having at least 2 active hydrogen groups relected from the group consisting of hydroxy, suthydryl, primary amino acids and secondary amino acids, one of said primary amino acids as accounting for one active hydrogen roup; and (c) a cohispocyanite.

The polyurethane resin according to claim 1 wherein said long-chain carboxylic acid is a C 36 dimer fathy acid.

- The polyurethane resin according to claim 1 wherein said carboxylic acid comprises between about 0 and 80 percent of said carboxylic acid component.
- The polyunathane resin according to claim 1 wherein said multi-functional compound is dimetrylolpropriedic acid.
 - A baseccat composition for deposition onto metal or plastic comprising:
 - (a) about 20 to 80% weight percent based on the final solids content of said basecost composition of an amonic polyurethane principal resin comprised of the reaction product of:
- (i) a polyester component with an alcohol having at least two hydroxyl groups wherein said carboxylic and component is comprised of at least about 50°s by weight of at least one long-chain carboxylic and having between 18 and 80 carbon atoms and at most about 50°s of at least one abor-chain dicarboxylic in
- (ii) a multi-functional compound having at least one active hydrogen and at least one carboxylic acid functionality:
- (iii) a compound having at least 2 active hydrogen groups selected from the group consisting of 15 hydroxyl, sulflydryl, primary amine, and secondary amine, one of said primary amines accounting for one active hydrogen and:
 - (iv) a polyisocyanate:(b) about 5 to about 50% by weight of an aminoplast cross-linking regin;
- (c) 0 to about 25 weight percent of a meciogy control agent selected from the group consisting of turned silica compaunds, benforite clays, and hectorite clays.
 - (d) 0 to ebout 35 weight percent of a branched chain polyester resin comprised of the reaction product of:

 (e) 0 to ebout 35 weight percent of a branched chain polyester resin comprised of the reaction product of:
- (i) a carboxylic acid component comprised of at least 50% by weight of at least one long chain carboxylic acid containing compound having between 18 and 60 carboxe, no more than about 48% by se weight of at least one short-chain diszaboxylic acid and between about 2 and 25% by weight of a polytuctional carboxylic acid or acid anilydrific, said polytuctional carboxylic acid or acid anthydrific.
 - having at least 3 carboxylic acid groups; and (ii) an alcohol component having an average functionality of at least 2 and
 - (e) about 2 to about 75 weight percent of a pigment-containing grind resin comprising:
- 30 (a) About 6 to about 60% by weight of said pigment-containing grind resin of a pigment: (b) About 20 to about 75% by weight of said pigment-containing grind resin of a polyurethane resin produced by the reaction product of:
- (i) a polyester resin component produced by the reaction of a carboxylic acid component comprised of at least 50% by weight of a long-chain carboxylic acid having between 18 and 60 carbon atoms and at most
- 39 about 50% of a short chain disarboxylic acid and an alcohol having at least 2 hydroxyl groups; and (ii) a mixture of a multi-functional compound having at least 1 active hydrogen and at least one carboxylic acid functionality, at least one compound having at least two active hydrogen groups, and a
- polyisocyanate, said carboxylic acid groups being neutralized with an amine; and
 (iii) c) About 20% to about 80% by weight of said pigment pastel pigment-containing grind resin of an
 aminoplast cross-linking agent.
 - The basecoat composition according to claim 5 wherein said polyurethane resin is comprised of C38 dimer fatty acid as the long-chain carboxylic acid.
 - The basecoat composition according to claim 5 wherein said polyfunctional carboxylic acid (d) is trimellitic anhydride.
- 45 8. The basecoat composition according to claim 5 wherein said hectorite clay is a purified sodium lithium magnesium sificate.
 - 9. A multi-layer peint composition comprising:
 - (a) A waterborne basecost composition comprising:
 - (i) an anionic polyurethane composition comprised of the reaction product of:
- (1) a polyester resin component produced by the reaction of a carboxylic acid component comprised of at least 50% by weight of least one long-thain carboxylic acid having between 18 and 90 carbon atoms and at most about 50% by weight of a short-chain dicarboxylic acid and an alcohol having at least 2 hydroxyl minister.
- (2) a mixture of at least one multi-functional compound having at least 1 active hydrogen group and at se least one carbonic acid functionality, at least one compound having at least two active hydrogen groups, and a polyspocyanide, suit carbonivic acid croups being neutralized with an amine.
 - (ii) a cross-linking agent;
 - (iii) a rheology control agent

(iv) optionally, a branched chain polyester resin comp prised of the reaction product of:

(1) a carboxylic acid component comprised of at least 50% by weight of a long-their carboxylic acid having between 18 and 90 carbons, no more bina about 46% by weight of a short-chain dicatorypic acid and between about 2 and 25% by weight of a polyfunctional carboxylic acid or acid anhydride having at a least 3 carboxylic acid crows: and

(2) an alcohol containing compound heving an average alcohol functionality of at least 2:(and)

(v) a pigment; and

(b) a clear topcost composition for overcoating said basecost composition.

10. The composition according to claim 9 wherein said long-chain carboxylic acid is C36 dimer fatty

A pigment-containing grind resin for use in waterborne basecoat compositions comprising:

(a) About 6 to about 60% by weight of said pigment-containing grind resin of a pigment;

(b) About 20 to about 75% by weight of said pigment-containing grind resin of a polyurethane resin

produced by the reaction product of:

(i) a polysetir resist component produced by the reaction of a carboxylic acid component comprised of
all test 50% by weight of a long-chain carboxylic acid flexing between 18 and 60 carbon atoms and at most

about 50% of a short chain dicarboxylic acid and an alcohol having at least 2 hydroxyl groups; and (i) a mixture of at least one multi-functional compound having at least it active hydrogen and at least one carboxylic acid functionality, et least one compound having at least two active hydrogen groups, and a

20 polyisocyanate, said carboxylic acid groups being neutralized with an amine; and (c) About 20% to about 80% by weight of said pigment-containing grind resin of an aminoplast cross-

linking agent.

12. A method of making a waterborne basecoat composition for use in a multi-layer coating comprising

the sequential staps of:

(a) Adding crosslinking agent in solution to a rheology control agent in solution and thoroughly mixino:

(b) Adding anionic polyurethane resin dispersion comprised of the reaction product of:

(i) a polyester resin component produced by the reaction of a carboxylic acid component comprised of a line set 50% by weight of a long-chain acid having between 18 and 60 carbon atoms and at most about 50% of a short-chain discrepositie acid and a stochet having at least 2 hydroxyl croust send

(ii) a mixture of a mutil-functional compound having at least 1 active hydrogen functionality and at least one carboxylic acid functionality, at least one compound having at least two active hydrogen groups, and a polylocyarate, said carboxylic acid group being neutralized with an amine, to the mixture of step (a) under adulation and mixing throughly.

(c) Adding a premixed sturry comprised of:

a sturry formed by premixing a branched chain polyester resin dispersion comprised of the reaction product of:

(1) a carboxylic acid component comprised of at least 50% by weight of a long chain carboxylic acid having between 15 and 60 carbons, no quere than about 46% by weight of a son-chain discharyoylic acid 49 and between about 2 and 25% by weight of a polytanctional carboxylic acid having at least 3 carboxylic acid crossis.

(2) an alcohol component heving an average functionality of at least 2

(3) water, and

(4) an amine-containing compound to the mixture obtained after step C;
(d) Anitating the mixture obtained from step B with the mixture from step C; and

(e) Adding Pigment Paste to the mixture obtained from step 0 and mixing thoroughly.

(f) Adjusting the pH and viscosity of the mixture obtained from step E.
13. The method according to claim 12 wherein said long-chain carboxylic acid is C38 dimer fatty acid.

14. A method of coating an automobile substrate with a multilayer coating comprising: (a) Applying to the primed substrate a first layer of waterborne coating composition comprised of:

(i) a polyurethane resin obtained form the reaction product of:

(i) a polyester resin component produced by the reaction of a carboxylic acid component comprised of
at least 50% by weight of a fonce-thain cathonytic acid hexing between 18 and 60 carbon atoms and at most

at least 50% by weight or a directional calculative action leaving desertine in earting or studies and a microabout 50% of a short chair disorboying and and an accord having at least 2 hydroxyl groups; and s (2) a mixture of a multi-functional compound having at least 1 active hydrogen functionality and at least one carboxylic paid functionality, at least one compound having at least two active hydrogen groups, and

polyisocyanate, said carboxylic acid groups being neutralized with an amine;

(ii) a cross-linking agent

- (iii) a rheology control agent; and
- (iv) a pigment paste;
- (v) optionally aluminum and/or mica (vii) optionally branched polyester resin
- (b) flash drying said first layer at a temperature of about room temperature and about 145 degrees F for between about 30 seconds and about 10 minutes; and
- (c) Applying at least one layer of a clear topcost onto said basecost: and
- (d) Curing said basecoat and topcoat to a hard, durable film.
- 15. The method according to claim 14 wherein said long-chain carboxylic acid is C36 dimer fatty acid. A multi-coated metal or plastic substrate comprising:
- A substrate coated with at least one waterborne basecost composition comprising:
- (a) about 20 to 80% weight percent based on the final solids content of said basecost composition of a first anionic polyurethane resin comprised of the reaction product of:
- (i) a polyester component with an alcohol having at least two hydroxyl groups wherein said carboxylic rs and component is comprised of at least about 50% by eight of a long-chain carboxylic acid having between 18 and 60 carbon atoms and at most about 50% of a short chain dicarboxylic acid:
 - (ii) a multi-functional compound having at east 1 active hydrogen functionality and at least one carboxylic acid functionality
 - (iii) a compound having at least 2 active hydrogen groups:
 - (iv) a polyisocyanate; and

acid.

- (v) en amine-containing compound for neutralizing the free carboxylic acid groups.
- (b) About 5 to about 50 weight percent of an minoplast cross-linking resin:
- (c) About 0.1 to about 25 weight percent of a rheology control agent, selected from the group consisting of furned silica component, bentonite clevs, and hectorite clavs:
- (d) About 0 to about 35 weight percent of a branched chain polyester resin comprised of the reaction product of:
- (i) a carboxylic acid component comprised of t least 50% by weight of a long chain carboxylic acid having between 18 and 60 carbons, no more than about 46% by weight of short-chain dicarboxylic acid and between about 2 and 5% by weight of a polyfunctional carboxylic acid or acid anhydride, said polyfunc-30 Itonal carboxylic acid having at leasst 3 carboxylic acid groups; and
 - (ii) an alcohol component having an average functionality of at least; and
- (e) About 2 to about 75 weight percent of a gigment, each of said basecost compositions being flashdried before being coated with 2) A clear topocating, said basecuat composition and said topocating being cured to a hard durable
- 17. The composition according to claim 16 wherein said long chain carboxylic acid is C36 dimer fatty